

(II) EP 1 193 306 A2

(12) EUROPEAN PATENT APPLICATION

(43) Date of publication: 03.04.2002 Bulletin 2002/14 (51) Int CI.7: **C09K 11/78**, C09K 11/79, C09K 11/66

- (21) Application number: 01123144.6
- (22) Date of filing: 27.09.2001
- (84) Designated Contracting States:

 AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU

 MC NL PT SE TR

 Designated Extension States:

 AL LT LV MK RO SI
- (30) Priority: 29.09.2000 JP 2000299320 09.03.2001 JP 2001066318
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- (54) Fluorescent substances for vacuum ultraviolet radiation excited light-emitting devices
- (57) A fluorescent substance for vacuum ultraviolet rediation soutce light-netting devices which enhalt light extract produces with enhalt light with a high luminance and is subject to only a small decrease in luminance due to exposure to plasma or the like, and a vacuum ultraviolet radiation excited light-emitting device including the fluorescent substance, are provided by year of a fluorescent substance for vacuum ultraviolet radiation excited light-emitting devices including: a compound represented by the general formular ImMO-nMG-2xMG-2xMceral Mis at least one met-

al selected from Ca. Sr and Ba, M² is at least one metal seselected from Mg and Zn, M³ is at least one metal seselected from Si and Ge, and m and n satisty 0.5 ≤ m≤3.5 and 0.5 ≤ m≤2.5. respectively, provided that when m=n=1. M¹ is either at least two metals selected from Ca, Sr and Ba or one of Sr and Ba; and at least one metal selected from Eu and Mn as an activator, and a vacuum ultraviolet radiation exclict-light-emitting devece including the fluorescent substance.

Description

[0001] The present invention relates to fluorescent substances suitably used for vacuum ultraviolet radiation excited light-emitting devices such as plasma display panels (hereinafter referred to as "PDPs") and rare ras

[0002] Fluorescent substances that emit light when excited with e.g. vacuum ultraviolet radiation include, for example, BaMgAl₁Q₇-Eu comprising Ba, Mg, Al, and 10 an activator (EU) as a blue fluorescent substance for vacuum ultraviolet radiation excited light-emitting devices, and Zn₂SiQ₂Mn comprising Zn, Si, O and an activator (Mn) as a green fluorescent substance. As well, (Y,Gd)BO₃:Eu comprising Y, Gd, B, O and an activator (EU), for example, has been known as a rod fluorescent substance. Such fluorescent substances are also used in rare gas lamps. These prior art fluorescent substance sore for vacuum ultraviolet radiation excited light-emitting devices, however, are desired to exhibit further en-panced luminance.

[0003] in vacuum ultraviolet radiation excited lightemitting devices such as PDPs and rare gas lamps, causing discharge in a rare gas generates plasma and then, vacuum ultraviolet radiation is generated. The prior aft fluorescent substances involve a problem that the luminance thereof decreases as they are exposed to plasma. Thus, there is a desire for a fluorescent substance for vacuum ultraviolet radiation excited lightemitting devices of which luminance does not much decrease due to exposure to plasma.

[0004] US-A-5.839,718 discloses a silicate fluorescent substance comprising a compound represented by the general formula: m(Sr_{1.0}M¹₀)O·n(Mg_{1.b}M²_b)O· 2 (Si_{1-c}Ge_c)O₂:Eu_xLn_v, wherein M¹ is Ba, M² is at least 35 one element selected from the group consisting of Be. Zn and Cd. Ln is at least one element selected from the group consisting of Sc. Y, La, Ce, Pr, Nd, Sm, Gd, Tb, Dv. Ho, Er, Tm, Yb, Lu, B, Al, Ga, In, Tl, Sb, Bi, As, P, Sn, Pb. Ti, Zr. Hf. V. Nb. Ta. Mo. W. Cr and Mn. Specific ex- 40 amples of such silicate fluorescent substances disclosed include Sr_{1.995}MgSi₂O₇:Eu_{0.005},Dy_{0.025},Cl_{0.025}, and Sr_{0.445}Ba_{1.55}MgSi₂O₇:Eu_{0.005},Dy_{0.025},Cl_{0.025}. These fluorescent substances are for use as light accumulators for display in dark places or for similar purposes. "Ex- 45 tended abstracts of the sixth international conference on the science and technology of display phosphors", pages 21-24, discloses CaMgSi₂O₆:Eu as a fluorescent substance for vacuum ultraviolet radiation excited lightemitting devices.

[0005] Such fluorescent substances comprising silicate still have the problem that their luminance is insufficient and decreases due to exposure to plasma.

[0006] It is an object of the present invention to provide a fluorescent substance for vacuum ultraviolet radiation excited light-emitting devices which emits light with a higher luminance and a small decrease in luminance after the exposure to plasma. [0007] This object could be achieved on the basis of the finding that, among silicate or germanate fluorescent substances for vacuum ultraviolet radiation excited fight-emitting devices, a fluorescent substance having a specific composition comprising: a compound represented by the general formula. mM*Io.nM*2-0.2M*O_{ch} wherein M* is at least one metal selected from Ca, Sr and Ba, M* is at least one metal selected from Mg and Cry, M* is at least one metal selected from Mg and Cry, M* is at least one metal selected from Mg and can man in satisfy 0.5s.ms.35 and 0.5s.ms.25 repoctively, provided that when m=1, M* is either at least two metals selected from Ca, Sr and Ba, or one of Sr and Ba, and at least one metal selected from Eu and Mn as an activator, emits light with a higher luminance and a smaller decrease in luminance after the exposure

to plasma. [0008] Accordingly, the present invention provides a fluorescent substance for vacuum ultraviolet radiation excited light-emitting devices, comprising; a compound represented by the general formula, mM1O·nM2O·2M3O₂, wherein M1 is at least one metal selected from Ca, Sr and Ba. M2 is at least one metal selected from Mg and Zn, M3 is at least one metal selected from Si and Ge, and m and n satisfy 0.5≦m≤3.5 and 0.5≦n≤2.5, respectively, provided that when m=n=1. M1 is either at least two metals selected from Ca. Sr and Ba. or one of Sr and Ba: and at least one metal selected from Eu and Mn as an activator. The present invention also provides a vacuum ultraviolet radiation excited light-emitting device comprising the fluorescent substance defined above.

[009] The fluorescent substance for vacuum ultraviolet radiation excited light-meitting device according to the present invention is characterized by comprising: a 5 base crystall formed of a compound represented by the general formule: mM10-mM2-0M2-0M2-, wherein MI is at least one metal selected from Ca, Sr and Ba, MF is least one metal selected from Ca, Sr and Ra and Is assist one metal selected from Si and Ge, and man in satisfy 0.5≤m35.8 and 0.5≤m25.8 people/wby provided that when m=n=1, MI is either at least two metals selected from Ca, Sr and Ba, or one of Sr and Ba, and at least noe metal selected from Eu and Mn as an advivation.

[0010] When m=n=1 in the above-noted general fortis mula, M¹ is selected from Sr, Ba, the combination of Sr, Ba and Ca, combination of Sr and Ca, combination of Ba and Sr, and combination of Ba and Ca.

[0011] Preferably, the fluorescent substance for vacuum ultraviolet radiation excited light-emitting devices according to the present invention has a crystal structure as same as diopside, akermanite, or merwinite [0012] Among fluorescent substances having the

same crystal structure as diopside, one comprising a compound of the above-noted general formula where 35 m=n=1 is proferable. A more preferable one as a fluorescent substance for vacuum ultraviolet radiation excited light-ientifliar devices is of a composition in which M1 and M2 are partially replaced with the activators Eu and Mn, respectively and which is represented by the general formula, M^1 = $E_{\rm u}/M^2$ = $M_{\rm u}/M_{\rm e}/M_{\rm o}/M^3$ $Q_{\rm e}/M_{\rm e}$ wherein M¹ is either at least two metals selected from Ca, Sr and Ba, or one of Sr and Ba, M² is at least one metal selected from Mg and Zn, M³ is at least one metal selected from Mg and Zn, M³ is at least one metal selected from S island Ce, and a and b satisfy $0 \le a \le 0.5$, $0 \le b \le 0.5$, and $0 \le a \le 0.5$.

[0013] A still preferred one of fluorescent substances having the same crystal structure as diopside and comprising a compound of the above-noted general formula where m=1 has a composition in which M* is Ca and S*, M* and M* and M* gand S*, respectively, and Ca is partially replaced with the activator Eu and which is represented by the general formular Cap—26F,EUM,MS*LO₆ wherein c and d satisfy 0<c≤0.1 and 0<d≤0.1, respectively.

[0014] Among fluorescent substances having the same crystal structure as akermanite, one comprising a compound of the above-noted general formula where m=2 and n=1 is preferable. A more preferable one as a fluorescent substance for vacuum ultraviolet radiation excited light-emitting devices has a composition in which M1 and M2 are partially replaced with the activators Eu and Mn, respectively and which is represented by the general formula: (M14...Eu_)a(M24.4Mn/)M3aOz. wherein M1 is at least one metal selected from Ca. Sr and Ba, M2 is at least one metal selected from Mg and Zn. M3 is at least one metal selected from Si and Ge. and e and f satisfy 0≤e ≤0.5, 0≤f≤0.5, and 0<e+f. Still more preferable is a fluorescent substance having a 30 composition in which f=0, and M3 is Si and which is represented by the general formula: (M14...Eu_)aM2 SiaO2. wherein M1 is at least one metal selected from Ca, Sr and Ba, M2 is at least one metal selected from Mg and Zn, and e satisfies 0.001≦e≦0.1.

[0015] Among fluorescent substances having the

same crystal structure as a kermanite, one comprising a compound of the above-noted general formula where m=1 and n=2 is preferable. A more preferable one as a fluorescent substance for vacuum ultraviolet radiation 40 excited light-emitting devices has a composition in which M1 and M2 are partially replaced with the activators Eu and Mn, respectively and which is represented by the general formula: (M11-hEuh)(M21-Mni)2M32O7, wherein M1 is at least one metal selected from Ca, Sr 45 and Ba, M2 is at least one metal selected from Mg and Zn. M3 is at least one metal selected from Si and Ge. and h and i satisfy 0≤h ≤0.5, 0≤i≤0.5, and 0<h+i. [0016] Among fluorescent substances having the same crystal structure as merwinite, one comprising a 50 compound of the above-noted general formula where m=3 and n=1 is preferable. A more preferable one as a fluorescent substance for vacuum ultraviolet radiation excited light-emitting devices has a composition in which M1 and M2 are partially replaced with the activators Eu and Mn, respectively and which is represented by the general formula: (M11-1Eu)3(M21-kMnk)M32O8, wherein M1 is at least one metal selected from Ca, Sr

and Ba, ${\rm M}^2$ is at least one metal selected from Mg and Zn, ${\rm M}^3$ is at least one metal selected from

Si and Ge, and j and k satisfy $0 \le j \le 0.5$, $0 \le k \le 0.5$, and 0 < j + k.

- [0017] Among fluorescent substances having the same crystal structure as diopside, akermanite or merwritlet, those having the same crystal structure as diopside and those having the same crystal structure as merwritle are preferable. Fluorescent substances having the same structure as diopside are particularly prefera-
- [0018] Raw materials usable as calcium, strontium and barium sources in the preparation of the fluorescent substance of the present invention may be either those which can be decomposed into xoide as thigh temperatures such as hydroxides, carbonates, fittrates, haides and oxalates of calcium, strontium and barium haigh piptip upity (99% or more), or oxides of celcium, strontium and barium haiden haiden such as the calcium, strontium and barium haiden haiden such as the calcium, strontium and barium haiden haiden such as the calcium, strontium and barium haiden haiden such as the calcium, strontium and barium haiden haiden such as the calcium, strontium and barium haiden haiden such as the calcium, strontium and barium haiden haiden such as the calcium strontium and barium haiden haiden such as the calcium strontium and barium haiden such as the calcium strontium and barium such as the calcium strontium stronti
- [0019] Raw materials usable as magnesium and zinc sources may be either those which can be decomposed into oxides at high temperatures such as hydroxides, carbonates, nitrates, halides and oxalates of magnesium and zinc having high purity (9% or oxides of magnesium and zinc having high purity (99% or more).

[0020] Raw materials usable as allicon and germanium sources may be either those which can be decomposed into oxides at high temperatures such as hydroxides, carbonates, nitrates, halides, and oxalates of silicon and germanium having high purity (99% or more), or oxides of silicon and germanium having high purity (99% or more).

[0021] Raw materials containing europlum or manganess that will serve as an activator may be either tosewhich can be decomposed into oxides at high temperatures such as hydroides, carbonates, intrates, helices and oxalates of europlum and manganese having high purity (99% or more), or oxides of europlum and manganese having high purity (99% or more).

[0022] There is no particular limitation to the method of preparing the fluorescent substance of the present invention. The fluorescent substance may be prepared, for example, by mixing the aforementioned raw materials and calcining the mixture.

[0023] For instance, the fluorescent substance having the composition represented by the formula: (M*_{1-E,M}), (M*_{2-E,M}, which is one of the preferable compositions described above, can be prepared by weighing and bending the foregoing aw materials into a predetermined composition, mixing them and calcining the mixture. In mixing such raw materials, it, is possible to use e.g., a ball mill. V-Type mixer or stirring device.

[0024] After the mixing, the mixture is calcined at a separature, for example, ranging between about 1000°C and about 1500°C for about 1 to about 100 hours, to give the fluorescent substance of the present invention. In the case where those raw materials which

can be decomposed into oxides at high temperatures such as hydroxides, carbonates, nitrates, halides and oxalates of the necessary elements are used, it is possible to conduct pre-calcination of the mixture at a temperature, for example, ranging between about 600°C 5 and about 900°C prior to the main-calcination.

[0025] There is no particular limitation to the catchining atmosphere but catchining in a reducing atmosphere comprising, for example, nitrogen or argon and hydrogen in an amount of 10% by othorne is preferable. The 10 pre-catchination may be performed either in atmospheric air or in a reducing atmosphere. To promote the calcining treaction, an appropriate amount of a flux may be added.

[0028] Further, the fluorescent substance obtained by 15 the above-described method may be pulverized using, for example, a ball mill or jet mill. Eurlher, the substance may be washed and classified. To obtain a fluorescent substance of higher crystallinity, re-calcination may be marked and classified.

[0027] The fluorescent substance of the present invention thus obtained is capable of emitting light with a high luminance when excited with vacuum ultraviolet radiation and exhibits only a small decrease in luminance due to exposure to plasma. In the manufacture of e.g. a PDP or rare gas lamp, a process typically includes dispersing the fluorescent substance in a solvent, adding a binder to the dispersion liquid, applying the dispersion liquid onto a light-emitting section, and heating the dispersion liquid thus applied to about 500°C to remove the 30 binder. The fluorescent substance of the present invention, even if subjected to such a process, exhibits only a small decrease in luminance due to the heat treatment. Accordingly, use of the fluorescent substance of the present invention in a vacuum ultraviolet radiation 35 excited light-emitting device such as PDP or rare gas lamp makes it possible to realize a PDP or rare gas lamp exhibiting a high luminance and a prolonged service life. Hence, the fluorescent substance of the present invention is suitable for vacuum ultraviolet radiation excited 40 light-emitting devices.

[0028] The fluorescent substance of the present invention can also be excited with ultraviolet radiation out of the vacuum ultraviolet region, x-rays or electron beam and hence may be employed in devices using ultraviolet radiation out of the vacuum ultraviolet region, X-rays or electron beam as an excitation source.

EXAMPLES

[0029] The present invention is in more detailed with reference to the following examples, but the invention is not limited to those examples.

EXAMPLE 1

[0030] Raw materials, namely, calcium carbonate (CaCO₂ produced by WAKOJUNYAKU CO., LTD.), stronium carbonate (SrCO₂ produced by WAKOUUN-YAKU CO. LTD., europium oxide (Eigu.G. produced by SHINETSU KAGAKUKOGYO CO. LTD.), basic magnesium carbonate (MpGO2_MgCOH_SH-Q.) produced by WAKOJUNYAKU CO., LTD.) and silicon oxide (SiO₂ produced by WAKOJUNYAKU CO., LTD.), were weighed and mixed together so that the molar ratio of CacO₂SrCO₃Eu₂O₃ ((MgCO₃)Mg(OH₂SH₂OSO₄ was 0.92150.0485, 0.0150.22. The resulting mixture was catclined at 1200°C for two hours in a stream of Argas containing H₂ in an amount of 2% by volume, the ground, and egain calcined at 1200°C for two hours in a stream of Arga containing H₂ in an amount of 2% volume. A fluorescent substance of a composition represented by the formula:

 $Ca_{0.275}Sr_{0.485}Eu_{0.00}MgSij_{Q_6} \ was thue obtained. When this fluorescent substance was irradiated with utwished tradiation using an exciment 148-mi lamp (Model: H0012 manufactured by USHIO DENKI CO) in a vacuum chamber under a pressure of 8.7 Pa (5 x 10^2 Torr) or lower, it emitted blue light with a luminance of 24 cd/m²$

[0031] The fluorescent substance thus obtained was subjected to a heat treatment at 500°C for 30 minutes in air. The fluorescent substance having undergone the heat treatment was measured for its luminance and, as a result, there was no decrease in luminance as compared with the luminance of the fluorescent substance before the heat treatment.

[0032] The fluorescent substance obtained was placed in a gas atmesphere having a X No composition ratio of 5% 55% by volume under a pressure of 13.2 Pa and exposed to plasma of 10 W for 50 minutes, two no plasma of 50 W for 15 minutes. After the exposure to plasma, the fluorescent substance was measured for its burniance and, as a result, there was no decrease in luminance as compared with the luminance of the fluorescent substance before the exposure to plasma.

[0033] The fluorescent substance obtained was subjected to a heat treatment at 500°C for 30 mutuses in air.
Subsequently, the fluorescent substance was placed in
a gas atmosphere having a Xe Ne composition ratio of
5% 95% by volume under a pressure of 1.2. Pe and
exposed to plasma of 10 W for 30 minutes, then to plasma of 50 W for 15 minutes. After the exposure to plasma,
the fluorescent substance was measured for its luminance and, as a result, there was a decrease as small
as 4% in fluminance as compared with the fuminance of
fluorescent substance before the heat treatment and
the exposure to plasma.

EXAMPLE 2

[0034] Raw materials, namely, strontium carbonate is (SrCO₃ produced by WAKOJUNYAKU CO, LTD), barium carbonate (BaCO₃ produced by WAKOJUNYAKU CO, LTD.), europium oxide (Eu₂O₃ produced by SHI-NETSU KAGAKUKOGYO CO., LTD.), basic magnesium carbonate ((MgCO₃)₄Mg(OH)₂-5H₂O produced by WAKOJUNYAKU CO., LTD.) and silicon oxide (SiO2 produced by WAKOJUNYAKU CO., LTD.), were weighed and mixed together so that the molar ratio of SrCO3.BaCO3:Eu2O3:(MgCO3)4Mg(OH)2-5H2O:SiO2 was 2.28:0.57:0.075:0.2:2. The resulting mixture was calcined at 1200°C for two hours in a stream of Ar gas containing H2 in an amount of 2% by volume. A fluorescent substance having a composition represented by the formula: Sr_{2.28}Ba_{0.57}Eu_{0.15}MgSi₂O₈ was thus obtained. When this fluorescent substance was irradiated with ultraviolet radiation using an excimer 146-nm lamp (Model: H0012 manufactured by USHIO DENKI CO.) in a vacuum chamber under a pressure of 6.7 Pa (5 x 10-2 Torr) or lower, it emitted blue light with a luminance of 15 30 cd/m².

EXAMPLE 3

[0035] Raw materials, namely, barium carbonate 20 EXAMPLE 5 (BaCO₂ produced by KANTOKAGAKU CO, LTD.), europium oxide (Eu₂O₃ produced by SHINETSU KAGAKU-KOGYO CO., LTD.), magnesium oxide (MgO produced by KANTOKAGAKU CO., LTD.) and silicon oxide (SiO₂ produced by KOJUNDOKAGAKU KENKYUSHO CO... LTD.), were weighed and mixed together so that the molar ratio of BaCO₃:Eu₂O₃:MgO:SiO₂ was 1.98:0.01:1:2. 0.1 mol of B2O2 as a flux was added relative to 1 mol of the product, and the resulting mixture was sufficiently wet-blended in acetone with mortar, followed by drying. The mixed raw material thus obtained was put into a stainless steel mold and then pressurized at 40 Mpa to form a circular pellet having a size of 15 mm (diameter) x 3 mm (thickness). The resulting pellet was put into an alumina crucible and calcined at 1200°C for three hours in an atmosphere having a H2 Ar composition ratio of 5% 95% by volume. A fluorescent substance having a composition represented by the formula: Ba_{1.98}Eu_{0.02}MgSi₂O₇ was thus obtained. When this fluorescent substance was irradiated with ultraviolet radi- 40 bluish green light with a high luminance. ation using an excimer 146-nm lamp (Model: H0012 manufactured by USHIO DENKI CO.) in a vacuum chamber under a pressure of 6.7 Pa (5 x 10-2 Torr) or lower, it emitted green light with a luminance of 95 cd/ m².

EXAMPLE 4

[0036] Used as starting materials for the preparation of (Sr_{0.99}Eu_{0.01})₂MgSi₂O₇ were strontium carbonate (SrCO3 produced by KANTOKAGAKU CO, LTD.), europium oxide (Eu₂O₃ produced by SHINETSU KAGAKU-KOGYO CO., LTD.), magnesium oxide (MgO produced by KANTOKAGAKU CO., LTD.) and silicon oxide (SiO₂ produced by KOJUNDOKAGAKU KENKYUSHO CO., LTD.). These materials were weighed and mixed together so that the molar ratio of SrCO3:Eu2O3:MgO:SiO2 was 1 98:0.01:1:2, then 0.1 mol of B2O3 as a flux was

added relative to 1 mol of the product, i.e., (SroopEun on)oMgSioO7, and the resulting mixture was sufficiently wet-blended in acetone with mortar, followed by drying. The mixed raw material thus obtained was put into a stainless steel mold and then pressurized at 40 Mpa to form a circular pellet having a size of 15 mm (diameter) x 3 mm (thickness). The resulting pellet was put into an alumina crucible and calcined at 1200°C for three hours in an atmosphere having a H2 Ar composition ratio of 5% 95%. When the sample resulting from the calcining was irradiated with ultraviolet radiation having a wavelength of 254 nm or 365 nm, it emitted light blue light with a high luminance in either case. When the sample was irradiated with ultraviolet radiation using an excimer 146-nm lamp (manufactured by USHIO DENKI CO.) in a vacuum chamber under a pressure of 6.7 Pa (5 x 10-2 Torr) or lower, it emitted intensive light blue light with a luminance of 25 cd/m2.

[0037] Used as starting materials for the preparation of (SrnogEunna)2ZnSi2O2 were strontium carbonate (SrCO₂), europium oxide (Eu₂O₂), zinc oxide (ZnO) and silicon oxide (SiO₂). These materials were weighed and mixed together so that the molar ratio of SrCO3:Eu2O3: ZnO:SiO2 was 1.98:0.01:1:2, then 0.1 mol of B2O3 as a flux was added relative to 1 mol of the product, i.e., (Sr_{0.99}Eu_{0.01})₂ZnSi₂O₇, and the resulting mixture was sufficiently wet-blended in acetone with mortar, followed by drying. The mixed raw material thus obtained was put into a stainless steel mold and then pressurized at 40 Mpa to form a circular pellet having a size of 15 mm (diameter) x 3 mm (thickness). The resulting pellet was put into an alumina crucible and calcined at 1200°C for three hours in an atmosphere having a H₂ Ar composition ratio of 5% 95%. When the sample resulting from the calcining was irradiated with ultraviolet radiation having a wavelength of 254 nm or 365 nm, it emitted

COMPARATIVE EXAMPLE 1

[0038] Raw materials, namely, calcium carbonate (CaCO3 produced by WAKOJUNYAKU CO. LTD.), europium oxide (Eu2O3 produced by SHINETSU KA-GAKUKOGYO CO., LTD.), basic magnesium carbonate ((MgCO₃)₄Mg(OH)₂·5H₂O produced by WAKOJUN-YAKU CO., LTD.) and silicon oxide (SiO2 produced by WAKOJUNYAKU CO., LTD.), were weighed and mixed together so that the molar ratio of CaCO3 Eu2O3: (MgCO₃)₄Mg(OH)₂-5H₂O:SiO₂ was 0.95:0.025:0.2:2. The resulting mixture was calcined at 1200°C for two hours in a stream of Ar gas containing Ho in an amount 55 of 2% by volume. A fluorescent substance having a composition represented by the formula: Can as Euro as MgSi2Os was thus obtained. When this fluorescent substance was irradiated with ultraviolet radiation using an excimer 146-nm lamp (Model: H0012 manufactured by USHIO DENKI CO.) in a vacuum chamber under a pressure of 6.7 Pa (5 x 10-2 Torr) or lower, it emitted blue light with a luminance of 12 cd/m2.

COMPARATIVE EXAMPLE 2

[0039] A commercially-available blue light emitting fluorescent substance (BaMgAl₁₀O₁₇:Eu) was subjected to a heat treatment at 500°C for 30 minutes in air. 10 The fluorescent substance having undergone the heat treatment was measured for its luminance and, as a result, there was a 1% decrease in luminance as compared with the luminance of the fluorescent substance before the heat treatment.

[0040] The commercially-available blue light emitting fluorescent substance (BaMgAl₄₆O₄₇:Eu) was placed in a gas atmosphere having a Xe Ne composition ratio of 5% 95% by volume under a pressure of 13.2 Pa and exposed to plasma of 10 W for 30 minutes, then to plas- 20 ma of 50 W for 15 minutes. After the exposure to plasma. the fluorescent substance was measured for its luminance and, as a result, there was a 25% decrease in luminance as compared with the luminance of the fluorescent substance before the exposure to plasma.

[0041] The commercially-available blue light emitting fluorescent substance (BaMgAl₁₀O₁₇:Eu) was subjected to a heat treatment at 500°C for 30 minutes in air. Subsequently, the fluorescent substance was placed in a gas atmosphere having a Xe Ne composition ratio of 30 5% 95% by volume under a pressure of 13.2 Pa and exposed to plasma of 10 W for 30 minutes, then to plasma of 50 W for 15 minutes. After the exposure to plasma. the fluorescent substance removed from the gas atmosthere was a 28% decrease in luminance as compared with the luminance of the fluorescent substance in a state before undergoing the heat treatment and the exposure to plasma.

[0042] The fluorescent substance of the present in- 40 vention emits light with a high luminance and exhibits only a small decrease in luminance due to exposure to plasma and, hence, is particularly suitable for use in vacuum ultraviolet radiation excited light-emitting devices such as PDPs and rare gas lamps. Since the fluorescent 45 substance makes it possible to realize vacuum ultraviolet radiation excited light-emitting devices exhibiting high luminance, it is highly useful for industry.

Claims

1. A fluorescent substance for vacuum ultraviolet radiation excited light-emitting devices comprising: a compound represented by the general formula. 55 mM1O-nM2O-2M3O2, wherein M1 is at least one metal selected from Ca, Sr and Ba, M2 is at least one metal selected from Mg and Zn, M3 is at least

one metal selected from Si and Ge, and m and n satisfy 0.5≤m≤3.5 and 0.5≤n≤2.5, respectively, provided that when m=n=1. M1 is either at least two metals selected from Ca. Sr and Ba. or one of Sr and Ba: and at least one metal selected from Euand Mn as an activator

- 2. The substance according to claim 1, which has the same crystal structure as diopside.
- 3. The substance according to claim 2, wherein m=n=1.
- 4. The substance according to claim 2, wherein M1 is Ca and Sr, and M2 and M3 are Mg and Si, respectively and the activator is Eu.
- 5. The substance according to claim 4, which is of a composition represented by the formula: Ca_{1-c-d}Sr_cEu_dMgSi₂O₆, wherein c and d 0<c≤0.1 and 0<d≤0.1, respectively.
- 6. The substance according to claim 1, which has the same crystal structure as akermanite.
- 7. The substance according to claim 6, wherein m=2 and n=1.
- 8. The substance according to claim 6, wherein M3 is SI and the activator is Eu.
- 9. The substance according to claim 6, wherein m=1 and n=2.
- phere was measured for its luminance and, as a result, 35 10. The substance according to claim 1, which has the same crystal structure as merwinite.
 - 11. The substance according to claim 10, wherein m=3 and n=1.
 - 12. A vacuum ultraviolet radiation excited light-emitting device comprising a fluorescent substance according to any one of claims 1 to 11.

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